9,10-DIHYDROXYETEARIC ACID

(low-melting thomer)

(Octadecanoic acid, 9,10-dihydroxy-)

CH₈(CH₂)₇CH=CH(CH₂)₇CO₂H + H₂O₂ + HCO₂H +
Otato acid

CH₈(CH₂)-CH-CH(CH₂)-CO₃H

CH₃(CH₂)₇CH—CH(CH₂)₇CO₂H + 2NaOH →

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CH₈(CH₂)₇CH—CH(CH₂)₇CO₂Na + HCO₂Na

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 $CH_3(CH_2)_7CH$ — $CH(CH_2)_7CO_2Na$ + HCI \rightarrow

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CH₈(CH₂)₇CH—CH(CH₂)₇CO₂H + NaCl OH OH

Low-melting isomer

Submitted by Daniel Swern, John T. Scanlan, and Geraldine B. Dickel.¹ Checked by John D. Roberts and Edgar F. Kiefer.

1. Procedure

To a well-stirred mixture of 141 g. (0.5 mole) of oleic acid (Note 1) and 425 ml. of formic acid (Note 2) in a 1-l. three-necked flask at 25° is added the appropriate amount (Note 3) of 30% (100 volume) hydrogen peroxide (approximately 60 g.) over a 15-minute period (Note 4). The reaction becomes mildly exothermic after a lag of about 5-10 minutes, and homogeneous after about 20-30 minutes. The temperature is maintained at 40° with a cold water bath at the beginning, and with a warm water bath or heating mantle toward the end, of the reaction.

fter about 3 hours or after analysis has indicated that the stide has been communed (Note 5), the formic acid is removed sy distillation under reduced pressure (b.p. 50°/125 mm.) in a tream of gas (carbon dioxide or nitrogen) to prevent bumping The residue in the fizsk, which consists of hydroxyiric acids, is heated for 1 hour at 100° with an excess ticous sodium bydroxide, and the hot, amber-colored in is cautiously poured into an excess of SN hydroa said with stirring. The oil which separates is allowed to be and the aqueous layer is discarded. The tan-colored the steam bath by addition of hot water and over califust sales and water-soluble acids soil has soldfilled, the squeous layer is disellitus proken into small pieces and dissolved in ethanol by beating on the steam bath. After of tox several invirus the product is collected on a secum. The yield of coule 9 10 diny mag 85-90° After a second 260 gg map: 83-90 Arts second 260 mil. of 95% ethanoù the product weighs 88s et about 90-92%. A third recrystallizay to produce a pure product melting at rsall yield is 55-60 g. (50-55%, based on the id) (Note 8).

2. Notes

The checkers used commercial v.s.r. oleic acid, which has a terme number of about 60-70 and contains 65-75% oleic term. The submitters report that, if highly purified oleic acid trace, the visit of fairly pure 9.10-dihydroxystearic acid is light contained that the purification procedure for oleic acid business. Preparations, 2, 100 (1952)] is more lengthy and inconvenient than the purification of the hydroxylation product. The over-all yield is approximately the same in either case.

The formic acid employed is the 98-100% grade. The sub-

The formic acid employed is the 98-100% grade. The submitters report that the 90% grade of acid is satisfactory, but the reaction mixture remains heterogeneous throughout. They also state that, instead of formic acid, an equal volume of glacial

acetic acid containing 2.5% by weight of concentrated sulfuric acid may be employed. With acetic acid-sulfuric acid, a 6-hour reaction time is required. However, the yield of 9,10-dihydroxy-stearic acid is slightly lower than the yield obtained when formit acid is employed and the lodine number of the crude reaction

product is about 6-9.

3. If commercial oleic acid is used, the todine number should be determined beforehand and the quantity of hydrogen parasish, adjusted accordingly. The hydrogen peroxide should be sampled immediately before use: "100 volume peroxide" usually tonsom about 30% bydrogen peroxide by weight. This determines a conveniently carried out by weighing 0.2-0.3 g. of the hydrogen peroxide solution into an Erlenmeyer flash with a greater stopper and adding 20 ml. of a glacial acetic acid elifornism solution (3:2 by volume). Two millitiers of saturated separate stand for 5 minutes. Distilled water (75 ml.) is added and disliberated iodine titrated with 0.1N sodium thiosuitate solution to a starch end point. This procedure is also satisfactory for determining the peroxide content of the oxidation mistate, except that 1-2 g. samples are taken [cf. Wheeler, Oil & Soop, 0, 89 (1932)].

4. The submitters state that in one-tenth scale preparations the hydrogen peroxide solution can be added in 1 portion. In larger

runs the addition may require 30 minutes to 1 hour.

5. The reaction time ranges from 1.5 to about 4 hours. Frogress of the reaction should be followed by determining the peroxide content of the oxidation mixture at half-hour intervals after all the hydrogen peroxide has been added. Approximately all the peroxide should be consumed before distillation is at-

tempted.

6. Instead of removing the formic acid by distillation, the reaction mixture may be poured into a large quantity of water and the oily layer dissolved in ether. The ether solution is washed free of formic acid and then subjected to distillation to remove the ether; hydroxyformoxystearic acids are left as a residue. The submitters found that, in larger-scale operations (five or more times the size of the run described), no ether was

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7. The pH of the wash water should be below 6 in order to be earlied that all scap in the product has been converted to free acid. If the pH is above 6, a small quantity of 3N hydrochloric acid should be added and the stirring continued for several minutes.

8. The submitters report that the high-melting isomer of 9.10 dihydroxystearic acid can be prepared from elaidic acid Biochem Preparations, 3, 118 (1953)] by essentially the procedure described for oleic acid. With elaidic acid, instead of removing the formic acid by distillation, the reaction mixture may be poured into hot water and the oil which forms separated mechanically. The product is not readily soluble in ether. When acetic acid containing sulfuric acid is employed as the solvent, the reaction mixture is poured into hot water with thorough mixing, allowed to cool to room temperature, and filtered. The subsequent procedure (saponification and acidification) is the same as that described for the hydroxylation of bleic acid except that the crude dihydroxystearic acid, obtained after acidification of the soap, cannot be melted with hot water during the washing but is merely stirred well at 95-100° on the steam bath with a large quantity of hot water (Note 7). About 5 ml. of ethanol per gram of solute should be used in the recrystallization. The pure product melts at 130-131°. The yield depends on the purity of the starting material; if highly purified elaidic acid is used, the yield is about 80% after one recrystallization.

3. Methods of Preparation

The procedures described have been published.² Other procedures, which are not so satisfactory as the ones described, have also been published.^{3,4}

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MWERINELLE AGAGICO SIEMON WROEK ANTHE

(Cyclohexanemethylamine; NAI-dimethyl-)

$$CO_2H$$

$$+ SOCl_2 \rightarrow \qquad + SO_3 + HCl$$

$$+ 2HN(CH_3)_2 \rightarrow \qquad + (CH_3)_2NH \cdot HCl$$

$$CON(CH_6)_2 \qquad + (CH_3)_2NH \cdot HCl$$

$$CON(CH_6)_2 \qquad CH_2N(CH_6)_2$$

Submitted by ARTHUR C. COPE and ENGELBERT CIGAREK. Checked by WILLIAM E. PARHAM and ROBERT KONCOS.

1. Procedure

A. N,N-Dimethylcyclohexanecarboxamide. In a 1-1. three-necked flask equipped with a reflux condenser and a dropping funnel, both carrying drying tubes, is placed 128 g. (1.0 mole) of cyclohexanecarboxylic acid (Note 1). Thionyl chloride (179 g., 1.5 moles) (Note 1) is added during 5 minutes to the acid, with stirring by a magnetic stirrer. The flask is placed in an oil bath and heated at a bath temperature of 150° for 1 hour. The reflux condenser is then replaced by a distillation head (Note 2), 200 ml. of anhydrous benzene is added, and the mixture is distilled until